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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/723,250	11/26/2003	Thomas M. DiMauro	3518.1024-000	6059
21005 75500 002266010 HAMILTON, BROOK, SMITH & REYNOLDS, P.C. 530 VIRGINIA ROAD P.O. BOX 9133 CONCORD, MA 01742-9133			EXAMINER	
			MAEWALL, SNIGDHA	
			ART UNIT	PAPER NUMBER
			1612	
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Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Application No. Applicant(s) 10/723 250 DIMAURO ET AL. Office Action Summary Examiner Art Unit Snigdha Maewall 1612 -- The MAILING DATE of this communication appears on the cover sheet with the correspondence address --Period for Reply A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) OR THIRTY (30) DAYS. WHICHEVER IS LONGER, FROM THE MAILING DATE OF THIS COMMUNICATION. Extensions of time may be available under the provisions of 37 CFR 1.136(a). In no event, however, may a reply be timely filed after SIX (6) MONTHS from the mailing date of this communication. If NO period for reply is specified above, the maximum statutory period will apply and will expire SIX (6) MONTHS from the mailing date of this communication - Failure to reply within the set or extended period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133). Any reply received by the Office later than three months after the mailing date of this communication, even if timely filed, may reduce any earned patent term adjustment. See 37 CFR 1.704(b). Status 1) Responsive to communication(s) filed on 20 October 2009. 2a) This action is FINAL. 2b) This action is non-final. 3) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213. Disposition of Claims 4)\ Claim(s) 1-5.8-10.21-25.27-30.60.70.89.91 and 92 is/are pending in the application. 4a) Of the above claim(s) is/are withdrawn from consideration. 5) Claim(s) _____ is/are allowed. 6) Claim(s) 1-5, 8-10, 21-25, 27-30, 60, 70, 89 and 91-92 is/are rejected. 7) Claim(s) _____ is/are objected to. 8) Claim(s) _____ are subject to restriction and/or election requirement. Application Papers 9) The specification is objected to by the Examiner. 10) The drawing(s) filed on is/are; a) accepted or b) objected to by the Examiner. Applicant may not request that any objection to the drawing(s) be held in abevance. See 37 CFR 1.85(a). Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d). 11) The oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152. Priority under 35 U.S.C. § 119 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: Certified copies of the priority documents have been received. 2. Certified copies of the priority documents have been received in Application No. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. Attachment(s)

U.S. Patent and Trademark Office PTOL-326 (Rev. 08-06)

1) Notice of References Cited (PTO-892)

Paper No(s)/Mail Date

Notice of Draftsperson's Patent Drawing Review (PTO-948)

3) Information Disclosure Statement(s) (PTO/SB/08)

Interview Summary (PTO-413)
 Paper No(s)/Mail Date.

6) Other:

5) Notice of Informal Patent Application

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DETAILED ACTION

 Receipt of Applicants' arguments /remarks filed on 10/20/09 is acknowledged.

Claims 6-7, 26 and 57-58 have been canceled. Claims 11-20, 31-56, 59, 61-69, 71-88 and 90 have been withdrawn.

Claim 60 has been amended.

Accordingly, claims 1-5, 8-10, 21-25, 27-30, 60, 70, 89 and 91-92 are being examined on the merits herein.

The following are new rejections which are not necessitated by claim amendments, according this office action is made non final. The previous rejections have been withdrawn in light of applicant's arguments.

Claim Rejections - 35 USC § 103

- The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
- (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.
- Claims 1-3, 5, 8-10, 21-23, 25, 27-30, 70, 89 and 91-92 are rejected under 35
 U.S.C. 103(a) as being unpatentable over Radomsky (US 5,942,499) in view of

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US PG pub. 20050054595 ('595).

Radomsky teaches administration of growth factors to bone. Radomsky teaches a bone growth-promoting composition comprising growth factors such as fibroblast growth factor and platelet-derived growth factor and their methods of use (column 1, lines 19, 35-36, and 61). The invention can be used in various sites of desired bone growth including vertebral compression fractures and in pathological bone defects associated with osteoporosis (column 2, lines 50 and 55-58). The invention describes an injectable mixture of growth factor for intraosseous, or within bone, administration (column 12, lines 5-12).

The reference does not teach administration of anti-resorptive agent, such as remicade.

'595 teach chondrocyte therapeutic system, title. In paragraph [0029], the reference teaches compounds which produce a therapeutic effect are remicade. Instant claims 91 and 92 teach antiresorptive agent to be remicade, therefore, the agent is considered to possess quality of being highly specific cytokine antagonist comprising a monoclonal antibody that inhibits TNF alfa as claimed.

"It is prima facie obvious to combine two compositions each of which is taught by the prior art to be useful for the same purpose, in order to form a third composition to be used for the very same purpose[T]he idea of combining them flows logically from their having been individually taught in the prior art." In re Kerkhoven, 626 F.2d 846, 850, 205 USPQ 1069, 1072 (CCPA 1980) (citations omitted).

In view of the facts recited above, it would have been prima facie obvious to a

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person of ordinary skill in the art at the time the invention was made to combine the prior art elements according to known methods to yield predictable results. The prior art teaches all of the limitations of the claimed invention. Radomsky teaches injectable mixture of growth factor for intraosseous or within bone administration of bone forming agent and '595 teaches administration of remicade for chondrocye development. Since radomsky teaches addition of mixtures, one would have been motivated to add remicade in addition to other bone forming agents and would have had reasonable expectation of success I obtaining an improves and better combination of treating osteoporosis or bone related treatment. The person of ordinary skill in the art would have been motivated to combine the references because both references teach combinatorial compositions comprising anti-TNF agents or osteoporosis treating agents. The person of ordinary skill would have reasonably expected success because the combinatorial compositions comprising bone forming agent and other active agents which help in treating osteoporosis were well known in the art at the time of the instant invention, as evidenced by the Radomsky et al. and one of skill in the art would have expected success by substituting the species of remicade as taught by the '595 patent. The person of ordinary skill in the art could have combined the elements as claimed by known methods to produce a composition comprising remicade and bone forming agents. One of skill in the art would have recognized that the results of the combination would have yielded nothing more than predictable results to one of ordinary skill in the art at the time the invention was made, as demonstrated by the teachings of the radomsky and '595 publication. Further, "It is prima facie obvious to combine two

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compositions each of which is taught by the prior art to be useful for the same purpose, in order to form a third composition to be used for the very same purpose ...[T]he idea of combining them flows logically from their having been individually taught in the prior art." In re Kerkhoven, 626 F.2d 846, 850, 205 USPQ 1069, 1072 (CCPA 1980) (citations' omitted). Since Radomsky teaches administration the bone forming agent to bone, it would have been obvious to one of ordinary to administer the bone forming agent and antiresorptive agent such as remicade to hip or any vertebral part of body. From the teachings of the reference, it is apparent that one of ordinary skill in the art would have had a reasonable expectation of success in producing the claimed invention. Therefore, the invention as a whole would have been prima facie obvious to one of ordinary skill in the art at the time the invention was made, as evidenced by the references, especially in the absence of evidence to the contrary.

 Claims 4 and 24 are rejected under 35 U.S.C. 103(a) as being unpatentable over Radomsky (US 5,942,499) in view of US PG pub. 20050054595 ('595) as discussed above and further in view of Boyle et al. (US 2003/0207827).

The references discussed above do not teach patient as post menopausal.

Boyle et al. teach methods to treat bone diseases such as osteoporosis comprising osteoprotegrin, which is a polypeptide that plays a role in promoting bone accumulation (page 1, paragraphs [0001] and [0006]). Boyle et al. further teach treatment of osteoporosis in postmenopausal women and a direct relationship between osteoporosis and incidence of hip and neck fractures (page 9, paragraph [0095]).

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Osteoprotegrin acts as a receptor of the TNF family and prevents receptor-ligand interaction (page 4, paragraph [0043]). Osteoprotegrin also blocks interleukin (IL)1-α and IL1-β produced hypercalcemia (page 40, paragraph [0344]). Boyle et al. also teaches that estrogen is a known anti-resorptive agent (page 41, paragraph [0355]).

It would have been obvious to one of ordinary skill in the art at the time of instant invention to utilize the bone forming agent and remicade to post menopausal women as a patient motivated by the teachings of Boyle et al. teaching the administration of antiresoptive agents for treatment of osteoporosis in post menopausal women.

Claim 60 is rejected under 35 U.S.C. 103(a) as being unpatentable over
 Radomsky (US 5,942,499) in view of US PG pub. 20050054595 ('595) as discussed above and further in view of Trieu et al. ((US 2002/0026244).

The references discussed above do not teach implants.

Trieu teaches methods of implanting nucleus pulposus implants (page 1, paragraph [0007]). The method involves removal of the natural nucleus pulposus of the intravertebral disc and implantation of the nucleus pulposus of the invention (page 10, paragraph [0109]). The nucleus pulposus implant of the invention may contain pharmacological agents used to treat osteoporosis including a bone morphogenetic protein, growth factors such as fibroblast growth factor and platelet-derived growth factor, and steroids (page 9, paragraphs [0101] and [0104]). The device of Trieu is placed adjacent to unfractured bones (page 9, paragraphs [0104]). Since the nucleus pulposus implant of the invention may contain pharmacological agents used to treat

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osteoporosis including a bone morphogenetic protein (see page 9, paragraph [0101]), it would be obvious that the device can be used to treat fractured bones such as a hip bone. Thus Trieu teaches local administration in between bones.

It would have been obvious to the one of ordinary skilled in the art at the time the invention was made to incorporate highly specific cytokine antagonist such as remicade as taught by '595 to the teachings of Trieu since the reference teaches advantage of the same in treating osteoporosis with bone morphogenetic protein etc. One skilled in the art would have been motivated to administer into the bone the formulation comprising the bone forming agent and remicade because Trieu et al. successfully teach local administration of implants/drug in between bones in order to treat osteoporosis.

Response to Arguments

- Applicant's arguments with respect to claims 1-5, 8-10, 21-25, 27-30, 60, 70, 89
 and 91-92 have been considered but are moot in view of the new ground(s) of rejection.
- Any inquiry concerning this communication or earlier communications from the
 examiner should be directed to Snigdha Maewall whose telephone number is (571)272-6197. The examiner can normally be reached on Monday to Friday; 8:30 a.m. to
 5:00 p.m. EST.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Frederick Krass can be reached on (571) 272-0580. The fax phone number for the organization where this application or proceeding is assigned is 571-273-0580.

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Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

/Snigdha Maewall/

Examiner, Art Unit 1612

/Gollamudi S Kishore/

Primary Examiner, Art Unit 1612